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 NEWS 3 JUL 28 EPFULL enhanced with additional legal status information from the epoline Register
 NEWS 4 JUL 28 IFICDB, IFIPAT, and IFIUDB reloaded with enhancements
 NEWS 5 JUL 28 STN Viewer performance improved
 NEWS 6 AUG 01 INPADOCDB and INPAFAMDB coverage enhanced
 NEWS 7 AUG 13 CA/Caplus enhanced with printed Chemical Abstracts page images from 1967-1998
 NEWS 8 AUG 15 CAOLD to be discontinued on December 31, 2008
 NEWS 9 AUG 15 Caplus currency for Korean patents enhanced
 NEWS 10 AUG 27 CAS definition of basic patents expanded to ensure comprehensive access to substance and sequence information
 NEWS 11 SEP 18 Support for STN Express, Versions 6.01 and earlier, to be discontinued
 NEWS 12 SEP 25 CA/Caplus current-awareness alert options enhanced to accommodate supplemental CAS indexing of exemplified prophetic substances
 NEWS 13 SEP 26 WPIDS, WPINDEX, and WPIX coverage of Chinese and and Korean patents enhanced
 NEWS 14 SEP 29 IFICLS enhanced with new super search field
 NEWS 15 SEP 29 EMBASE and EMBAL enhanced with new search and display fields
 NEWS 16 SEP 30 CAS patent coverage enhanced to include exemplified prophetic substances identified in new Japanese-language patents
 NEWS 17 OCT 07 EPFULL enhanced with full implementation of EPC2000
 NEWS 18 OCT 07 Multiple databases enhanced for more flexible patent number searching
 NEWS 19 OCT 22 Current-awareness alert (SDI) setup and editing enhanced
 NEWS 20 OCT 22 WPIDS, WPINDEX, and WPIX enhanced with Canadian PCT Applications
 NEWS 21 OCT 24 CHEMLIST enhanced with intermediate list of pre-registered REACH substances

 NEWS EXPRESS JUNE 27 08 CURRENT WINDOWS VERSION IS V8.3,
 AND CURRENT DISCOVER FILE IS DATED 23 JUNE 2008.

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FILE COVERS 1907 - 18 Nov 2008 VOL 149 ISS 21
FILE LAST UPDATED: 17 Nov 2008 (20081117/EP)

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<http://www.cas.org/legal/info/policy.html>

```

=> s 6-methylmercaptopurine (w) riboside
 4190652 6
 326 METHYLMERCAPTOPURINE
 285 6-METHYLMERCAPTOPURINE
          (6 (W)METHYLMERCAPTOPURINE)
 4426 RIBOSIDE
 111 6-METHYLMERCAPTOPURINE (W) RIBOSIDE

```

=> s 11 and N-phosphonacetyl-L-aspartic acid
3243595 N
371 PHOSPHONACETYL
1711911 L
76309 ASPARTIC
4710213 ACID
62 N-PHOSPHONACETYL-L-ASPARTIC ACID
(N (W) PHOSPHONACETYL (W) L (W) ASPARTIC (W) ACID)
L2 3 L1 AND N-PHOSPHONACETYL-L-ASPARTIC ACID

=> s 12 and alanosine
 201 ALANOSINE
 L3 0 L2 AND ALANOSINE

=> s 12 and 3-bromopyruvate
 7433128 3
 1057 BROMOPYRUVATE
 150 3-BROMOPYRUVATE
 (3(W)BROMOPYRUVATE)
 L4 0 L2 AND 3-BROMOPYRUVATE

=> s 12 and adriamycin
 12068 ADRIAMYCIN
 L5 2 L2 AND ADRIAMYCIN

=> d 12 ibib abs hitstr 1-2

L2 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2008 ACS on STN
 ACCESSION NUMBER: 1996:125542 CAPLUS
 DOCUMENT NUMBER: 124:219696
 ORIGINAL REFERENCE NO.: 124:40305a,40308a
 TITLE: Enhanced antitumor activity of an adriamycin +
 5-fluorouracil combination when preceded by
 biochemical modulation
 AUTHOR(S): Stolfi, Robert L.; Colofiore, Joseph R.; Nord, L. D.;
 Martin, Daniel S.
 CORPORATE SOURCE: Catholic Medical Center, Woodhaven, NY, 11421, USA
 SOURCE: Anti-Cancer Drugs (1996), 7(1), 100-4
 CODEN: ANTDEV; ISSN: 0959-4973
 PUBLISHER: Rapid Science Publishers
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 AB A three-drug combination, PMA, consisting of (phosphonacetyl)-L-aspartic
 acid + 6-methylmercaptopurine riboside +
 5-aminonicotinamide, preceding either 5-fluorouracil (5-FU) or adriamycin
 (Adr), produced tumor-regressing activity in a murine advanced breast
 tumor model not attainable with either 5-FU or Adr as single agents, or
 with any lesser combination of these drugs administered at maximally
 tolerated doses. Marked tumor-regressing activity was further increased
 significantly by using 5-FU and Adr together in conjunction with the
 modulatory biochemical conditioning (particularly ATP depletion) provided by
 pretreatment with PMA.

L2 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2008 ACS on STN
 ACCESSION NUMBER: 1994:95764 CAPLUS
 DOCUMENT NUMBER: 120:955764
 ORIGINAL REFERENCE NO.: 120:16835a,16838a
 TITLE: Chemotherapeutic drug combinations
 INVENTOR(S): Martin, Daniel S.; Stolfi, Robert L.; Colofiore,
 Joseph R.; Nord, L. D.
 PATENT ASSIGNEE(S): USA
 SOURCE: PCT Int. Appl., 79 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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WO 9323014	A1	19931125	WO 1993-US4775	19930520
W: AU, CA, JP				
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
AU 9343834	A	19931213	AU 1993-43834	19930520
AU 684709	B2	19980108		
EP 641193	A1	19950308	EP 1993-914010	19930520
R: AT, BE, CH, DE, FR, GB, IE, IT, LI, NL				
JP 08506317	T	19960709	JP 1993-503842	19930520
PRIORITY APPLN. INFO.:			US 1992-885809	A 19920520
			WO 1993-US4775	A 19930520

AB Drug combinations for the treatment of neoplastic diseases comprise (1) cellular energy depletion compns. containing an inhibitor of purine nucleotide biosynthesis, a nicotinamide antagonist, and optionally an inhibitor of pyrimidine nucleotide biosynthesis and (2) apoptosis-inducing agents. For example, antineoplastic effects of combinations of N-(phosphonacetyl)-L-aspartic acid, 6-methylmercaptopurine riboside, 6-aminonicotinamide, and 5-fluorouracil in breast tumor-bearing mice were demonstrated.

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FILE 'CAPLUS' ENTERED AT 17:04:46 ON 18 NOV 2008

L1	111 S 6-METHYLMERCAPTOPURINE (W) RIBOSIDE
L2	3 S L1 AND N-PHOSPHONACETYL-L-ASPARTIC ACID
L3	0 S L2 AND ALANOSINE
L4	0 S L2 AND 3-BROMOPYRUVATE
L5	2 S L2 AND ADRIAMYCIN

=> d 12 ibib abs hitstr 3

L2 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2008 ACS on STN
 ACCESSION NUMBER: 1992:542977 CAPLUS
 DOCUMENT NUMBER: 117:142977
 ORIGINAL REFERENCE NO.: 117:24565a,24568a
 TITLE: Biochemical modulation of tumor cell energy:
 regression of advanced spontaneous murine breast
 tumors with 5-fluorouracil-containing drug combination
 AUTHOR(S): Stolfi, Robert L.; Colofiore, Joseph R.; Nord, L. D.;
 Koutcher, Jason A.; Martin, Daniel S.
 CORPORATE SOURCE: Cancer Res. Dep., Cathol. Med. Cent., Woodhaven, NY,
 11421, USA
 SOURCE: Cancer Research (1992), 52(15), 4074-81
 CODEN: CNREA8; ISSN: 0008-5472
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 AB This report describes a highly active chemotherapeutic drug combination, consisting of N-(phosphonacetyl)-L-aspartate plus 6-methylmercaptopurine riboside plus 6-aminonicotinamide plus 5-fluorouracil, in CD8F1 mice bearing spontaneous, autochthonous, breast tumors or first-passage advanced transplants of these spontaneous tumors. The combination and sequence of administration of these drugs were selected on the basis of known potentiating biochem. interactions. High performance liquid chromatog. and NMR spectroscopy measurements of biochem. changes resulting from treatment with N-(phosphonacetyl)-L-aspartate plus 6-methylmercaptopurine riboside plus 6-aminonicotinamide indicated a severe depletion of cellular energy levels in the treated

tumors. 6-Aminonicotinamide produced a severe block of the pentose shunt, and 5-fluorouracil severely inhibited both thymidylate synthase and thymidine kinase in the treated tumors. This quadrupole drug combination, administered on a 10-11-day schedule, produced an impressive partial tumor regression rate of 67% of large, spontaneous, autochthonous, murine breast tumors and a tumor regression rate of 74% of first-passage transplants of the spontaneous breast tumors.

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L5 2 S L2 AND ADRIAMYCIN

=> d 15 ibib abs hitstr 1-2

L5 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1996:125542 CAPLUS

DOCUMENT NUMBER: 124:219696

ORIGINAL REFERENCE NO.: 124:40305a,40308a

TITLE: Enhanced antitumor activity of an adriamycin + 5-fluorouracil combination when preceded by biochemical modulation

AUTHOR(S): Stolfi, Robert L.; Colofiore, Joseph R.; Nord, L. D.; Martin, Daniel S.

CORPORATE SOURCE: Catholic Medical Center, Woodhaven, NY, 11421, USA

SOURCE: Anti-Cancer Drugs (1996), 7(1), 100-4

CODEN: ANTDEV; ISSN: 0959-4973

PUBLISHER: Rapid Science Publishers

DOCUMENT TYPE: Journal

LANGUAGE: English

AB A three-drug combination, PMA, consisting of (phosphonacetyl)-L-aspartic acid + 6-methylmercaptopurine riboside + 5-aminonicotinamide, preceding either 5-fluorouracil (5-FU) or adriamycin (Adr), produced tumor-regressing activity in a murine advanced breast tumor model not attainable with either 5-FU or Adr as single agents, or with any lesser combination of these drugs administered at maximally tolerated doses. Marked tumor-regressing activity was further increased significantly by using 5-FU and Adr together in conjunction with the modulatory biochem. conditioning (particularly ATP depletion) provided by pretreatment with PMA.

L5 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1994:95764 CAPLUS

DOCUMENT NUMBER: 120:95764

ORIGINAL REFERENCE NO.: 120:16835a,16838a

TITLE: Chemotherapeutic drug combinations

INVENTOR(S): Martin, Daniel S.; Stolfi, Robert L.; Colofiore, Joseph R.; Nord, L. D.

PATENT ASSIGNEE(S): USA

SOURCE: PCT Int. Appl., 79 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

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WO 9323014	A1	19931125	WO 1993-US4775	19930520
W: AU, CA, JP				
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE				
AU 9343834	A	19931213	AU 1993-43834	19930520
AU 684709	B2	19980108		
EP 641193	A1	19950308	EP 1993-914010	19930520
R: AT, BE, CH, DE, FR, GB, IE, IT, LI, NL				
JP 08506317	T	19960709	JP 1993-503842	19930520
PRIORITY APPLN. INFO.:			US 1992-885809	A 19920520
			WO 1993-US4775	A 19930520
AB	Drug combinations for the treatment of neoplastic diseases comprise (1) cellular energy depletion compns. containing an inhibitor of purine nucleotide biosynthesis, a nicotinamide antagonist, and optionally an inhibitor of pyrimidine nucleotide biosynthesis and (2) apoptosis-inducing agents. For example, antineoplastic effects of combinations of N-(phosphonacetyl)-L-aspartic acid, 6-methylmercaptopurine riboside, 6-aminonicotinamide, and 5-fluorouracil in breast tumor-bearing mice were demonstrated.			

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